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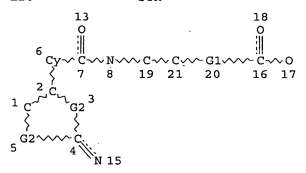
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STEREO ATTRIBUTES: NONE L18 STR

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STEREO ATTRIBUTES: NONE L20 STR



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GRAPH ATTRIBUTES: RSPEC I NUMBER OF NODES IS 16

STEREO ATTRIBUTES: NONE

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L24 ANSWER 1 OF 7 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:878167 HCAPLUS

DOCUMENT NUMBER:

141:366227

TITLE:

Preparation of imidazolidin-2-one and oxazolidin-2-one derivatives as glucagon receptor antagonists/inverse

agonists

INVENTOR(S):

Kurukulasuriya, Ravi; Link, James T.; Patel, Jyoti R.;

Sorensen, Bryan K.

PATENT ASSIGNEE(S):

SOURCE:

U.S. Pat. Appl. Publ., 24 pp.

CODEN: USXXCO

DOCUMENT TYPE:

LANGUAGE:

Patent

USA

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 2004209928 A1 20041021 US 2003-743954 20031223 PRIORITY APPLN. INFO.: US 2002-437132P P 20021230

OTHER SOURCE(S):

GI

MARPAT 141:366227

I

AB Compds. of formula (I) or pharmaceutically suitable salts, esters or prodrugs thereof, [wherein A = CO2H, tetrazole; B = H, F, OH, alkoxy, NRaRb (wherein Ra, Rb = H, alkyl, alkylcarbonyl, alkylsulfonyl alkoxyalkyl, cycloalkyl, cycloalkylcarbonyl, cycloalkylsulfonyl, cycloalkylalkyl, heterocyclyl, heterocyclylalkyl, heterocyclylcarbonyl, heterocyclesulfonyl); D = aryl, heteroaryl; E = (CH2)n; m, n = 0, 1, 2; V = C(Rc), N (wherein Rc = H, alkyl, alkoxy, alkoxyalkyl, cycloalkyl, cycloalkyloxy, cycloalkylalkyl, heterocyclyl, heterocyclylalkyl); W = C(RdRe), (Rd)N, O, S, S(O), S(O)2; X = C(O), C(O)C(RfRg), C(RfRg)C(O), C(S), C(RfRg), C(RfRg), C(RiRj), C:N(Rj), S(O), S(O)2; Y = C(RkRm), (Rk)N, O, S, S(0), S(0)2; Z = a bond, C(RpRq), C(RpRq)C(RsRt); Rd, Re, Rf, Rg, Ri, Rj, Rk, Rm, Rp, Rq, Rs, Rt = H, alkyl, alkoxy, alkoxyalkyl, aryl, arylalkyl, aryloxy, arylalkoxy, cycloalkyl, cycloalkylalkyl, cycloalkyloxy, cycloalkylalkyl, heterocyclyl, heterocyclylalkyl, heterocyclyloxy, heterocyclylalkoxy] are prepared These compds. are novel glucagon receptor antagonists or inverse agonists and are useful for treating (1) type 2 diabetes in a mammal, (2) symptoms related to type 1 or type 2 diabetes in a mammal wherein said symptoms are selected from the group consisting of hyperglycemia, hyperinsulinemia, inadequate glucose clearance, obesity, hyperlipidemia, lipid metabolism disorders and hypertension, and (3) diabetes or syndrome X in a mammal. Thus, 3-[4-[1-(4-tert-butylcyclohexylamino)-2-(tertbutyldimethylsilanyloxy)ethyl]benzoylamino]propionic acid Et ester underwent addition reaction with 4-(trifluoromethoxy) phenyl isocyanate in THF at ambient temperature for 12 h to give 3-[4-[1-[N-(4-tert-butylcyclohexyl)-N'-(4-trifluoromethoxyphenyl)ureido]-2-(tert-butyldimethylsilanyloxy)ethyl]be nzoylamino)propionic acid Et ester (II). Desilylation of II with Bu4NF in THF at 0° for 30 min gave 3-[4-[1-[N-(4-tert-Butylcyclohexyl)-N'-(4-

trifluoromethoxyphenyl)ureido] -2-hydroxyethyl]benzoylamino]propionic acid
Et ester which was cyclized by treatment with polymer supported
triphenylphosphine (0.146 g, 0.44 mmol) followed by di-Et
azodicarboxylate, saponification with NaOH in aqueous MeOH, and acidification
with 1 N
 aqueous HCl to give N-[4-[3-(4-tert-butylcyclohexyl)-2-oxo-1-[4(trifluoromethoxy)phenyl]imidazolidin-4-yl]benzoyl]-β-alanine. The

compds. I were found to inhibit glucagon-stimulated cAMP production at a concentration of 20 μM a range of about 50 to .apprx.100%.

780763-68-0P, N-[4-[(2Z)-3-(4-tert-Butylcyclohexyl)-2-[[4-(trifluoromethoxy)phenyl]imino]-1,3-oxazolidin-4-yl]benzoyl]-β-alanine 780763-69-1P, N-[4-[(2Z)-2-[(4-Bromophenyl)imino]-3-(4-tert-butylcyclohexyl)-1,3-oxazolidin-4-yl]benzoyl]-β-alanine 780763-70-4P, N-[4-[(2Z)-3-(4-tert-Butylcyclohexyl)-2-[(4-phenoxyphenyl)imino]-1,3-oxazolidin-4-yl]benzoyl]-β-alanine 780763-71-5P, N-[4-[(2Z)-2-(1,1'-Biphenyl-4-ylimino)-3-(4-tert-butylcyclohexyl)-1,3-oxazolidin-4-yl]benzoyl]-β-alanine RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)
(preparation of imidazolidin-2-one and oxazolidin-2-one derivs. as glucagon receptor antagonists/inverse agonists for treating type II diabetes or symptoms related to type 1 or 2 diabetes)

RN 780763-68-0 HCAPLUS

Double bond geometry as shown.

RN 780763-69-1 HCAPLUS

CN β-Alanine, N-[4-[(2Z)-2-[(4-bromophenyl)imino]-3-[4-(1,1-dimethylethyl)cyclohexyl]-4-oxazolidinyl]benzoyl]- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 780763-70-4 HCAPLUS

CN β-Alanine, N-[4-[(2Z)-3-[4-(1,1-dimethylethyl)cyclohexyl]-2-[(4-phenoxyphenyl)imino]-4-oxazolidinyl]benzoyl]- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 780763-71-5 HCAPLUS

CN β-Alanine, N-[4-[(2Z)-2-([1,1'-biphenyl]-4-ylimino)-3-[4-(1,1-dimethylethyl)cyclohexyl]-4-oxazolidinyl]benzoyl]- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L24 ANSWER 2 OF 7 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:412749 HCAPLUS

DOCUMENT NUMBER: 140:423705

TITLE: A preparation of piperidinylcyclopentyl amide

derivatives, useful as modulators of chemokine

receptor activity

INVENTOR(S): Zhou, Changyou; Pasternak, Alexander; Yang, Lihu

PATENT ASSIGNEE(S): Merck & Co., Inc., USA SOURCE: PCT Int. Appl., 100 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.					KIND DATE			APPLICATION NO.						DATE			
	WO	2004	0411	63		A2		2004	0521	1	WO 2	003-1	JS34	099		2	003/10	024
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			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	OM,
			PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ТJ,	TM,	TN,
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PRIORITY APPLN. INFO.:								1	US 2	002-	4223	81P		P 20	0021	030		
OTHER GI	S SC	URCE	(S):			MAR	PAT	140:4	4237	05								

# \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

The invention relates to piperidinylcyclopentyl amide derivs. of formula I [wherein: X is -O-, -CH2O-, -CO2-, or -OC(O)-, etc.; W is (un)substituted Ph or heterocycle; Z is C, N, or O, wherein when Z is N, then R4 is absent, and when W is O, then both R3 and R4 are absent; n = 0-4; R1 is H, halo, trifluoromethyl, OH, alkyl, or CN, etc.; R2 is (un)substituted C0-6alkyl-(phenyl/heterocycle); R3 is (un)substituted C0-6alkyl-phenyl; R4 is H, OH, CN, or alkyl, etc.; R5 and R6 are independently selected from H, OH, alkyl, alkoxy, or oxo, etc.; R3 and R5 or R4 and R6 may be joined together to form (un)substituted ring], useful as modulators of chemokine receptor activity. In particular, these compds. are useful as modulators of the chemokine receptor CCR-2. For instance, piperidinylcyclopentyl amide derivative II (CCR-2 receptor binding IC50 < 1μM) was prepared via amination of the obtained intermediate cyclopentanone derivative III by 4-(4-fluorophenyl)piperidine with a yield of 66% (example 1).

IT 690654-26-3P 690654-27-4P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of piperidinylcyclopentyl amide derivs., useful as modulators of chemokine receptor activity)  $\,$ 

RN 690654-26-3 HCAPLUS

CN Glycine, N-[[(1R,3S)-1-(2-amino-4-thiazolyl)-3-(4-phenyl-1-piperidinyl)cyclopentyl]carbonyl]-N-[[3,5-bis(trifluoromethyl)phenyl]methy

1]-, ethyl ester, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

$$H_2N$$
 $R$ 

RN 690654-27-4 HCAPLUS

CN Glycine, N-[[(1R,3S)-1-[2-(acetylamino)-4-thiazolyl]-3-(4-phenyl-1-piperidinyl)cyclopentyl]carbonyl]-N-[[3,5-bis(trifluoromethyl)phenyl]methyl]-, ethyl ester, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

L24 ANSWER 3 OF 7 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2003:242293 HCAPLUS

DOCUMENT NUMBER: 138:271976

TITLE: Preparation of amino acid sulfonamide derivatives as

protease inhibitors

INVENTOR(S): Palmer, James

PATENT ASSIGNEE(S): Axys Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 100 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

P	ATENT :	NO.			KIN	D :	DATE			APPL	ICAT	ION 1	. 01		D	ATE	
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WC	2003	0249	23		<b>A1</b>		2003	0327	1	WO 2	002-	US28	505		2	0020	909
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		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	ΝZ,	PL,	PT,	RO,
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US	2003	1582	31		A1		2003	0821	1	US 2	002-	2375	09		2	0020	909
RI	ry App	LN.	INFO	. :					1	US 2	001-	3222	20P	:	P 2	0010	914
R S	SOURCE	(S):			MAR	PAT	138:	2719	76								

PRIOR OTHER

Sulfonamide compds. R6NHCR4R5CONHCHR3CH2SO2NR1R2 [R1, R2 = H, .AB (functionalized) alkyl, (hetero)cycloalkyl, (hetero)aryl, (hetero)bicycloaryl, functional groups, etc.; R3 = H, (functionalized) alkyl, etc.; R4 = H, alkyl; R5 = (functionalized) alkyl; or CR4R5 = cycloalkylene; R6 = H, acyl] or their pharmaceutically-acceptable salts were prepared as cysteine protease inhibitors. Thus, benzyl 1S-[[1S-[[(4-methoxyphenyl)sulfamoyl]methyl]-3-phenylpropyl]carbamoyl]-3methylbutylcarbamate was prepared by coupling of 2S-(benzyloxycarbonylamino)-4-methylpentanoic acid with 2S-amino-N-(4-methoxyphenyl)-4-phenylbutane-1sulfonamide hydrochloride in THF in the presence of 4-methylmorpholine and iso-Bu chloroformate.

294622-71-2 IT

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of amino acid sulfonamide derivs. as protease inhibitors)

294622-71-2 HCAPLUS RN

L-Leucine, N-[4-[2-(3-pyridinylamino)-4-thiazolyl]benzoyl]-, CNmonohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

HCl

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L24 ANSWER 4 OF 7 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2001:693319 HCAPLUS

DOCUMENT NUMBER:

135:257468

TITLE:

Preparation of N-(4-thiazolylbenzoyl)-N-(cyanomethyl)-

L-leucinamides and analogs as protease inhibitors

INVENTOR(S):

Palmer, James T.; Setti, Eduardo L.; Tian, Zong-Qiang;

Ι

Venkatraman, Shankar; Wang, Dan-Xiong

PATENT ASSIGNEE(S):

Axys Pharmaceuticals, Inc., USA

SOURCE:

PCT Int. Appl., 73 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	PATENT NO.					KIND DATE				APPLICATION NO.					DATE		
WO	WO 2001068645				A2 20010920			WO 2001-US8332						20010314			
WO	2001	0686	45		A3 20020307								•				
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PRIORITY	RIORITY APPLN. INFO.:							US 2000-189694P									
GI																	

AΒ The title compds. and their pharmaceutically acceptable salts, N-oxides, prodrugs, protected derivs., or isomers thereof were prepared as cysteine protease inhibitors. For example, stirring a solution of 4-[2-(1-tert-butoxycarbonylpiperidin-4-ylamino)thiazol-4-yl]benzoic acid (preparation given) and the MeSO3H salt of 2S-amino-N-cyanomethyl-4methylpentanamide overnight at room temperature with PyBOP and diisopropylethylamine in DMF, followed by conversion to the Et ester, yielded I (77%). Test compds. inhibited cathepsin B, K, L, and S (no data). The invention compds. and compns. with a bisphosphonic acid and/or an estrogen receptor agonist are claimed for treating osteoporosis in post-menopausal women (no data).

IT 294622-48-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of N-thiazolylbenzoyl-N-cyanomethyl-Lleucinamides and analogs as cysteine protease inhibitors for treatment of osteoporosis)

294622-48-3 HCAPLUS RN

CN L-Leucine, N-[4-[2-(3-pyridinylamino)-4-thiazolyl]benzoyl]-,
1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L24 ANSWER 5 OF 7 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:666701 HCAPLUS

DOCUMENT NUMBER: 133:252050

TITLE: Preparation of novel N-cyanomethyl amide compounds and

compositions as protease inhibitors to treat

osteoporosis

INVENTOR(S): Bryant, Clifford M.; Palmer, James T.; Rydzewski,

Robert M.; Setti, Eduardo L.; Tian, Zong-Qiang;

Venkatraman, Shankar; Wang, Dan-Xiong

PATENT ASSIGNEE(S): Axys Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 155 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

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WO 2000055126							WO 2000-US6837						20000315		
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OTHER SOURCE(S): MARPAT 133:252050

Title compds. [R1R2NCR3R4CN; R1 = R11R7NCR5R9X1, R11R8NCR6R10X2NR7CR5R9CX1; X1, X2 independently = CO, CH2SO2; R5, R6 independently = H, C1-6alkyl; R7, R8 independently = H, C1-6alkyl; R9, R10 independently = (un) substituted-C1-6alky1; R9-R7 = trimethylene, tetramethylene, phenylene-1,2-dimethylene; R10-R8 = trimethylene, tetramethylene, phenylene-1,2-dimethylene; R5-R9 = C3-8cycloalkylene, C3-8heterocycloalkylene; R10-R6 = C3-8cycloalkylene, C3-8heterocycloalkylene; R11 = X4X5R18; X4 = CO, COCO, SO2; X5 = bond, O, NH; R18 = C1-6alkyl; R2 = H, C1-6alkyl; R3 = H, C1-6alkyl; R4 = CN, COOH, COOC1-6alkyl; R2-R4 = trimethylene, tetramethylene, phenylene-1,2-dimethylene; R4-R3 = C3-8cycloalkylene, C3-8heterocycloalkylene], N-oxide, prodrug, isomers, pharmaceutically acceptable salts, and composition are prepared as therapeutically effective estrogen receptor agonist. Title compds. are claimed in treating osteoporosis in post-menopausal woman in which cathepsin K activity contributes to the pathol. and symptomatol. of the disease. Thus, the title compound (S)-C6H5CH2OCONHCH(CH2CH(CH3)2)CONHCH2CN was prepared

IT 294622-48-3P 294622-71-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of novel N-cyanomethyl amides and compns. as protease inhibitors)

RN 294622-48-3 HCAPLUS

CN L-Leucine, N-[4-[2-(3-pyridinylamino)-4-thiazolyl]benzoyl]-,
 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 294622-71-2 HCAPLUS

CN L-Leucine, N-[4-[2-(3-pyridinylamino)-4-thiazolyl]benzoyl]-, . monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

#### HC1

L24 ANSWER 6 OF 7 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2000:351518 HCAPLUS

DOCUMENT NUMBER:

133:4650

TITLE:

Preparation of heteroaryl-substituted aromatic

compounds as antiherpes compounds

INVENTOR (S):

Simoneau, Bruno; Crute, James J.; Faucher, Anne-Marie; Grygon, Christine A.; Hargrave, Karl D.; Thavonekham,

P 19981112

Bounkham

PATENT ASSIGNEE(S):

Boehringer Ingelheim (Canada) Ltd., Can.

SOURCE:

PCT Int. Appl., 157 pp.

DOCUMENT TYPE:

CODEN: PIXXD2 Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000029399	Al	20000525	WO 1999-CA1066	19991109

W: CA, JP, MX, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE

PRIORITY APPLN. INFO.: US 1998-108272P

OTHER SOURCE(S):

MARPAT 133:4650

GI

AB The title compds. X-Aryl-Y-Z [I; X = 5-6 membered aromatic heterocycle; Aryl = (un) substituted Ph, pyridyl; Y is absent or a bridging group, for example NHC(0)CH2; Z is a terminal group, for example NHCO2t-Bu or II], which inhibit the herpes helicase-primase enzyme, rendering the compds. useful as antiviral agents, were prepared E.g., a multi-step synthesis of benzamide III was presented. Biol. data (IC50 and/or EC50 against HSV-1 and HCMV) for compds. I were given.

IT 270566-77-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of heteroaryl-substituted aromatic compds. as antiherpes compds.)

RN 270566-77-3 HCAPLUS

CN L-Phenylalanine, N-[4-(2-amino-4-thiazolyl)benzoyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L24 ANSWER 7 OF 7 HCAPLUS COPYRIGHT 2005 ACS on STN

5

ACCESSION NUMBER:

1999:457919 HCAPLUS

DOCUMENT NUMBER:

131:116229

TITLE:

Preparation of thiazolecarboxamides as vitronectin

receptor antagonists

INVENTOR(S):

Alig, Leo; Edenhofer, Albrecht; Hilpert, Kurt; Weller,

Thomas

PATENT ASSIGNEE(S):

F. Hoffmann-La Roche AG, Switz.

SOURCE:

Eur. Pat. Appl., 87 pp. CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
			EP 1998-124670	
EP 928790	B1	20030305		
			GB, GR, IT, LI, LU, NL,	SE. MC. PT.
TR ST I.T	LV FT	PΩ		
US 6100282	A	20000808	US 1998-218567 NZ 1998-333590 NZ 1998-333591	19981222
NZ 333590	A	20000526	NZ 1998-333590	19981224
NZ 333591	A	20000526	NZ 1998-333591	19981224
AT 233746	E	20030315	AT 1998-124670	19981224
PT 928790	T	20030731	PT 1998-124670	19981224
ES 2193471	тз	20031101	AT 1998-124670 PT 1998-124670 ES 1998-124670	19981224
NO 9806159	Α	19990705	NO 1998-6159	19981228
		20020311		
ZA 9811925	Α	20000629	ZA 1998-11925	19981229
IL 127785				19981229
CA 2257328	AA	19990702	CA 1998-2257328	19981230
AU 9896144	A1	19990722	AU 1998-96144	19981230
AU 720618	B2	20000608		
SG 74686 JP 2000053664	A1	20000822	SG 1998-5978	19981230
JP 2000053664	A2	20000222	JP 1999-10	19990104
JP 3113237	B2	20001127		
BR 9900006	Α	20000411		19990104
MX 9900215 RU 2218337	A:	20000630		19990104
RU 2218337	C2	20031210	RU 1999-100277	19990105
HK 1020953			HK 1999-106136	19991228
US 6320054	, <b>B1</b>	20011120	US 2000-526033	20000315
US 2002010316	A1	20020124	US 2001-878704	20010611
US 6344562	B2	20020205		
PRIORITY APPLN. INFO.:			EP 1998-100006	
			US 1998-218567	A3 19981222
			US 2000-526033	A3 20000315

OTHER SOURCE(S): MARPAT 131:116229

R1 (CH2) aZ (CONR9) cZ1 (CH2) e (NB) fAm (NH) g (CH2) n [CH [ (CO) k (NH) 1R10] ] i (CH2) jCO2H[I; A = CO or SO2; B,R9 = H or (cyclo)alkyl; R1 = NR6CONR5(CH2)bR4, NR5R6, NHC(:NR8)NHR7, etc.; R4 = H, (cyclo)alkyl, (hetero)aryl; R5,R6 = H, (cyclo)alkyl, aryl, etc.; R7,R8 = H, (ar)alkyl, etc.; R7R8 = atoms to complete a ring; R10 = H, OH, (ar)alkyl, carboxy(alkyl), alkoxycarbonyl, etc.; Z = (un)substituted thiazole-2,4- or -2,5-diyl; Z1 = bond or arylene; a,j = 0-2; b = 0-4; c,f,g,h,i,k,l,m = 0 or 1; e = 0-3; h = 0-5] were prepared Thus, H2NC(:NH)NHCSNH2 was cyclocondensed with BrCH2COCO2Et and the saponified product amidated by H2NCH2CH2CONHCH2CH2CO2Et to give, after saponification, H2NC(:NH)NHZ(CONHCH2CH2)2CO2H (Z = thiazole-2,4-diyl).

Data

for biol. activity of I were given.

232593-43-0P 232593-44-1P 232593-45-2P IT

232593-46-3P 232593-60-1P 232593-89-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of thiazolecarboxamides as vitronectin receptor antagonists)

RN 232593-43-0 HCAPLUS CN Benzenepropanoic acid

Benzenepropanoic acid,  $\beta$ -[[4-[2-[(aminoiminomethyl)amino]-4-methyl-5-thiazolyl]benzoyl]amino]methyl]-4-chloro- (9CI) (CA INDEX NAME)

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PAGE 2-A

Cl

RN 232593-44-1 HCAPLUS

CN Benzenepropanoic acid,  $\beta$ -[[[3-[2-[(aminoiminomethyl)amino]-4-methyl-5-thiazolyl]benzoyl]amino]methyl]-4-chloro- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{NH} \\ & \text{H}_2\text{N}-\text{C}-\text{NH} \\ & \text{N} \\ & \text$$

RN 232593-45-2 HCAPLUS

CN Benzenepropanoic acid,  $\beta$ -[[4-[2-[(aminoiminomethyl)amino]-4-methyl-5-thiazolyl]benzoyl]amino]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{NH} \\ \parallel \\ \text{H}_2\text{N-C-NH} \\ \text{S} \\ \\ \text{C-NH-CH-CH}_2\text{-CO}_2\text{H} \\ \parallel \\ \text{O} \\ \text{Ph} \\ \end{array}$$

RN 232593-46-3 HCAPLUS

CN Benzenepropanoic acid,  $\beta$ -[[3-[2-[(aminoiminomethyl)amino]-4-methyl-5-thiazolyl]benzoyl]amino]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{NH} \\ & \parallel \\ & \parallel \\ & \text{H}_2\text{N-C-NH} \\ & \text{N} \end{array} \\ \text{HO}_2\text{C-CH}_2 \\ - \text{CH-NH-C} \\ & \parallel \\ & \text{Ph} \end{array}$$

RN 232593-60-1 HCAPLUS

CN Benzenepropanoic acid,  $\beta$ -[[[3-[2-[(aminoiminomethyl)amino]-4-thiazolyl]benzoyl]amino]methyl]-4-chloro- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{NH} & \text{O} & \text{CH}_2-\text{CO}_2\text{H} \\ \text{H}_2\text{N}-\text{C}-\text{NH} & \text{C}-\text{NH}-\text{CH}_2-\text{CH} \\ & \text{S} & & \text{C} \end{array}$$

232593-89-4 HCAPLUS RN

Butanoic acid, 4-[[3-[2-[(aminoiminomethyl)amino]-4-methyl-5-CN thiazolyl]benzoyl]amino]-2-[(butylsulfonyl)amino]-, monohydrochloride, (2S) - (9CI) (CA INDEX NAME)

### Absolute stereochemistry.

#### HCl

IT 232595-25-4P 232595-26-5P 232595-27-6P

232595-28-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of thiazolecarboxamides as vitronectin receptor antagonists)

RN 232595-25-4 HCAPLUS

CNBenzenepropanoic acid,  $\beta$ -[[[4-[2-[(aminoiminomethyl)amino]-4-methyl-5thiazolyl]benzoyl]amino]methyl]-4-chloro-, ethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

| Cl

RN 232595-26-5 HCAPLUS

CN Benzenepropanoic acid, β-[[[3-[2-[(aminoiminomethyl)amino]-4-methyl-5thiazolyl]benzoyl]amino]methyl]-4-chloro-, ethyl ester (9CI) (CA INDEX
NAME)

$$\begin{array}{c|c} & \text{NH} \\ & \text{H}_2\text{N}-\text{C}-\text{NH} & \text{N} \\ & \text{S} & \\ & \text{EtO-C-CH}_2 \\ & \text{CH-CH}_2-\text{NH-C} \\ & \text{C} \\ & \text{C} \\ \end{array}$$

RN 232595-27-6 HCAPLUS

CN Benzenepropanoic acid,  $\beta$ -[[4-[2-[(aminoiminomethyl)amino]-4-methyl-5-thiazolyl]benzoyl]amino]-, ethyl ester (9CI) (CA INDEX NAME)

RN 232595-28-7 HCAPLUS

CN Benzenepropanoic acid,  $\beta$ -[[3-[2-[(aminoiminomethyl)amino]-4-methyl-5-thiazolyl]benzoyl]amino]-, ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{NH} \\ \text{H}_2\text{N-C-NH} \\ \text{S} \\ \\ \text{Eto-C-CH}_2\text{-CH-NH-C} \\ \\ \text{Ph} \\ \text{O} \\ \end{array}$$

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REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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